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PATEMI Docker No. 514162000120

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In the application of

Che-Kum James SHEN

Sedal No. 10/014/220

Filing Date: November 9, 2001

For:

HS-40 ENHANCER-CONTAINING VECTORS IN TRANSCENIC ANIMALS

Examiner 3. Koushal Group Art Units 1633

DECLARATION OF CHE-KUNJAMES CHEN PURSUANT TO 17 C.F.R. 9 1.132

Assistant Commissioner for Palents Washington D.C. 20231

Dear Sh:

I Che-Kun James Shan, declare as follows:

- Lam currently employed as a Distinguished Research Fellow and Director at the Institute of Molecular Biology, Academia Sinica.
- I am the inventor of the invention disclosed in the above referenced parent application, and am familiar with the contents thereof. I have rasigned my rights in the invention to the Academia Simica and stand to receive 20% of profits in connection with the invention pursuant to my employment with Academia Sinica.
- I received a Ph.D. in Chemistry Bons the University of California, Besteley, July, 1977, and have been extively involved in molecular bidiogy and biotechnology-related research for 30 years. My corriculum vitus is satisfied lieuto as Exhibit A.

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- I am a co-author of Zhang et al. (JBC 279() 5):8501-8503, 1995) and therefore I am familiar with the contents thereof. In addition, I have resd the Office Action flored October 19, 2005 where the Examiner discussed this article. The translection assays that we conducted were transfert transfection assays. In a transfert transfection assay, the UNA construct does not integrate into the host cell's genome. This is particularly true in mammalian cells such as human cells because the random integration frequency in mammalian cells is very low under the conditions used for the translent transfection assay, in the range of one event per 18 18 cells (Roth, D. R. and Wilson, J. H. p. 621-651, Genetic Recombination, Am. Soc. Microbiol. 1988; Further, the tiene after transless transfection till assaying is too short (-48 hr) to allow the random integration to occur. Thus, few if any cell would have the construct integrated into the genome during transient transfection. Even when conditions are optimized to promote integration, the efficiency is still quite low. To oversome this low efficiency, according attempting to achieve integration of a vector use a selectable marker to kill cells that do not have the vector integrated into their senome. We did not perform any such selection stop for the paper. In addition, the vector used for transfert transfection did not have a selectable marker cay it that could have been used to select for integration.
- The TCTGAGTCA sequence provides the unexpected characteristic of position independent expression when integrated into the genome. Position independence can only be demonstrated when an expression construct is integrated into the genome of the host cell, not during translept expression assays. Therefore, position independent expression was not seen in our experiments for the Zhang et al. pages and would not have been presided from the results that we published in Zimny et al. One skilled in the art would have not predicted that this sequence provides position independent expression until results our patent application and the results therein.

January / 2005

2

Serial No. 10/014/229 Decide No. 514162000120

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